ARKHIPOV, Nikolay Nikolayevich; KARPACHEV. Pavel Spiridonovich:

MAYZEL', Maks Mikhaylovich, doktor tekhn. nauk, prof.;

PLEVAKO, Nikolay Alekseyevich; ZAYO CHKOVSKIY, A.B., doktor
tekhn. nauk, prof., retsenzent; ZCLOTOV, V.I., inzh., retsenzent; ZYBIR, V.P., doktor tekhn. nauk, retsenzent; KAPUSTIN,
I.I., doktor tekhn. nauk, prof., retsenzent; KOZLOV, B.A.,
inzh., retsenzent; POPOV, S.M., doktor tekhn. nauk, prof.,
retsenzent; EPPEL', S.S., kand. tekhn.nauk, dots., retsenzent; MINAYEVA, T.M., red.; SHVETSOV, S.V., tekhn. red.

[Basic processes, machinery, and apparatus of light industry]
Osnovnye protsessy, mashiny i aparaty legkoi promyshlemosti.
[By] N.N.Arkhipov i dr. Moskva, Izd-vo nauchno-tekhn. lit-ry
RSFSR, 1961. 491 p. (MIRA 15:2)

ARKHIPOV, N.N.; KARPACHEV, P.S.; MAYZEL', M.M., doktor tekhn. nauk, prof.; PLEVAKO, N.A.; UDODOVSKIY, A.N.; kand. tekhn. nauk, retsenzent; RYZHOVA, L.P., red. izd-va; EL'KIND, V.D., tekhn. red.

[Fundamentals of the design and construction of standard machines and devices for light industry] Osnovy konstruirovaniia i rascheta tipovykh mashin i apparatov legkoi promyshlennosti. [By] N.N.Arkhipov i dr. Pod red. M.M. Maizelia. Moskva, Mashgiz, 1963. 599 p. (MIRA 16:7) (Machinery-Design and construction) (Instruments)

Whatereva, A.A., kand. tekhn. nak, dotsent; W. Lenkova, I..., doktor tekhn. nak, prof.; CHETHOV, H.V., dok or tekhn. navi., prof.; KARPACHEV, P.S., inch.

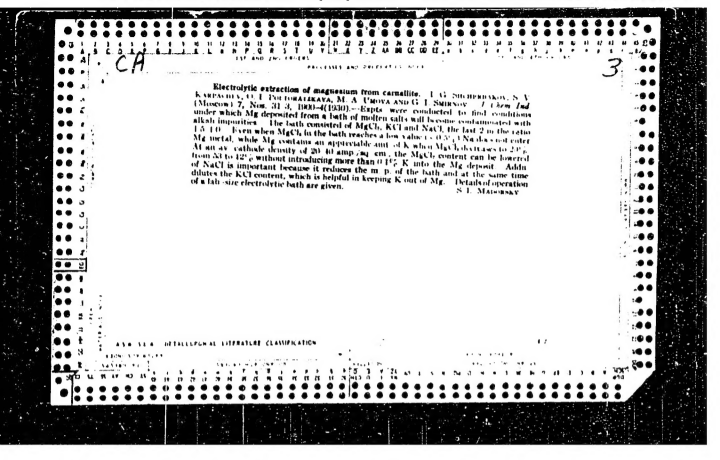
Effect of mechanical actions on the agest ration of the penetration in tanning tanning. Nauch. trudy MTIAS is 0.003-93-93.

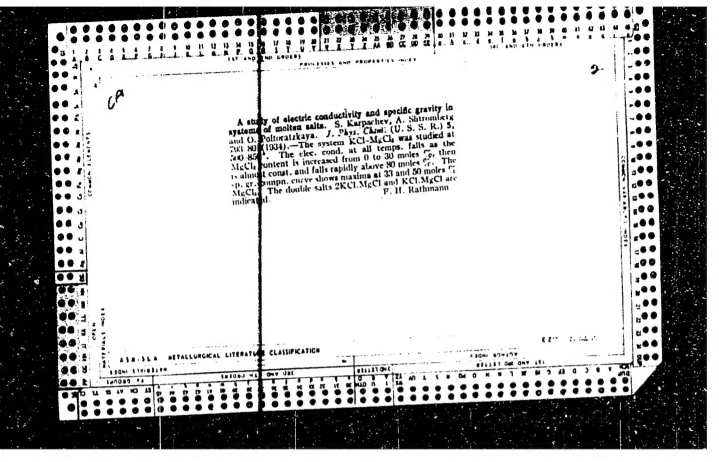
(NIRA 17:11)

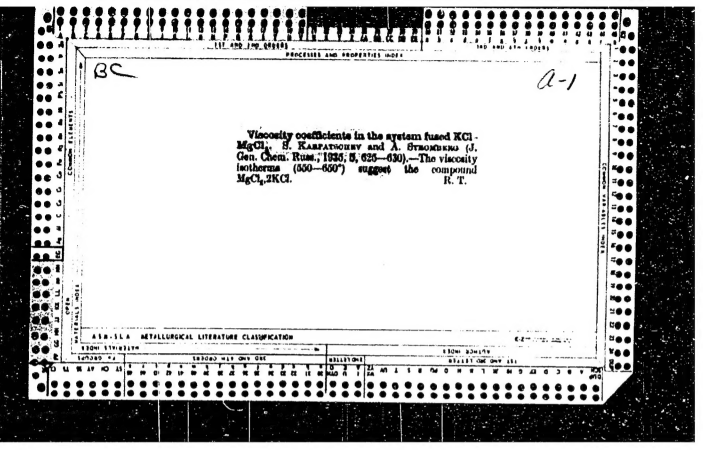
l. Karedra tekhnologii kozhi i mekha la  $\rightarrow$ skogo tek sallogichoshogo instituta legkov promyshlennosti.

KARPACHEV, Pavel Spiridonovich; MAYZEL, Maks Mikhaylovich, doktor tekhn.nauk.prof.; FLEVAKO, Nikolay Alekseyevich; CHETKIN, Fetr Petrovich; ZAYONCHKOVSKIY, A.D., doktor tekhn.nauk.prof., retsenzent; ZOLOTOV, V.I., inzh., retsenzent

[Machinery and apparatus for the manufacture of artificial leather and film materials] Lashiny i apparaty prolayedsty Iskuastvennol kezhi i plenochnykh materialov. [By] P.S.Karpachev i dr. Moskya, Legkala industriia, 1964. 609 p. (MIRA 18:2)

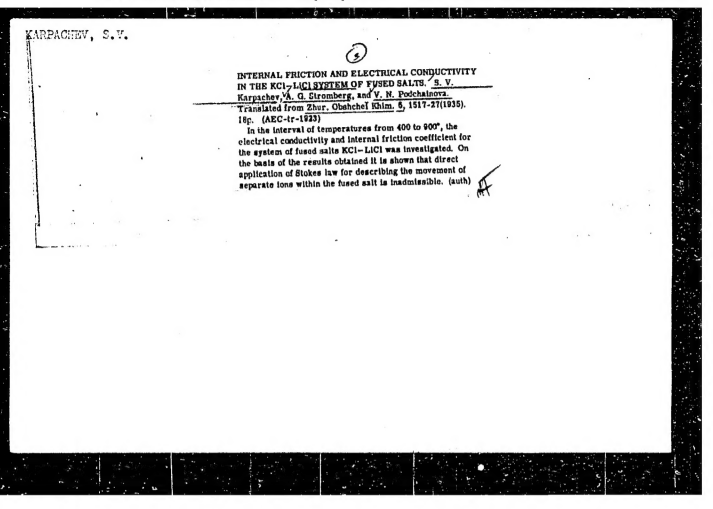


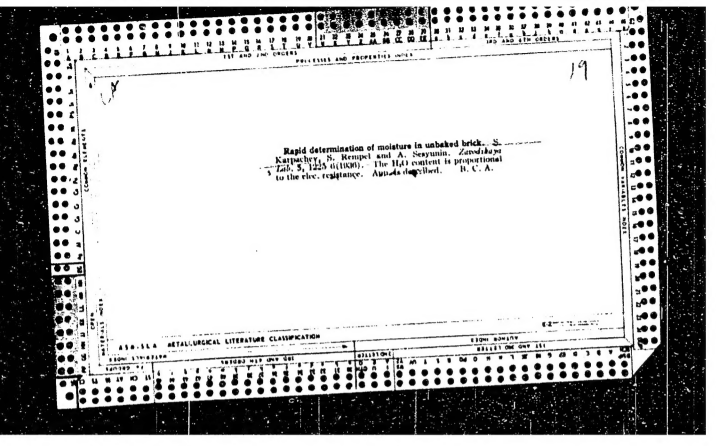


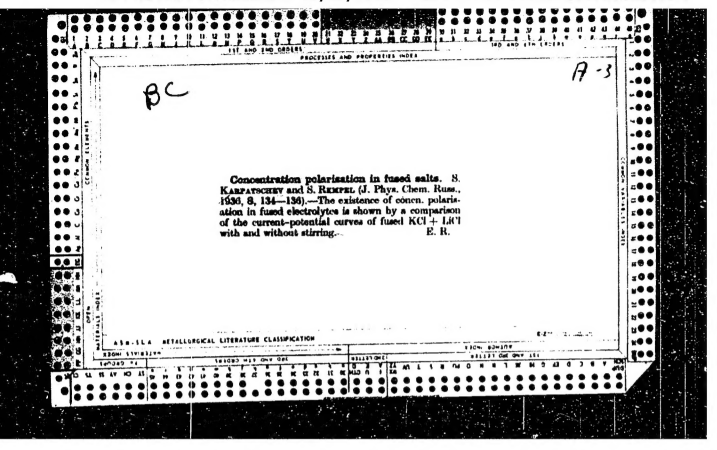


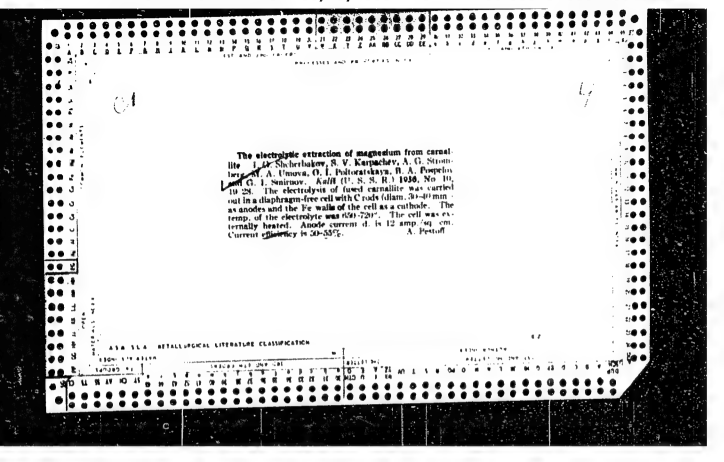
### "APPROVED FOR RELEASE: 06/13/2000 CI

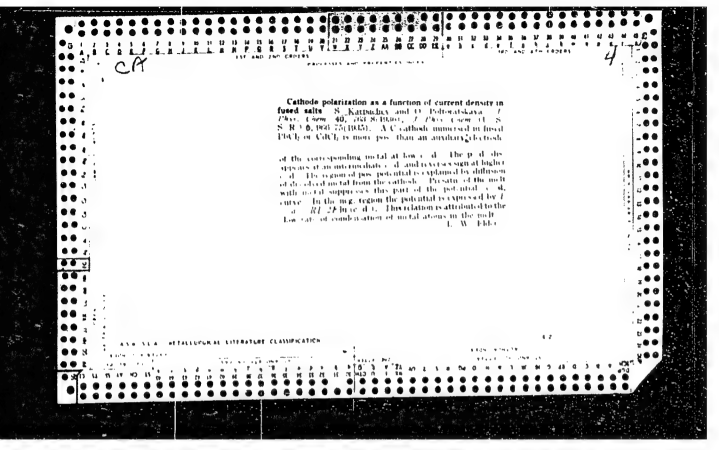
CIA-RDP86-00513R000720820003-9

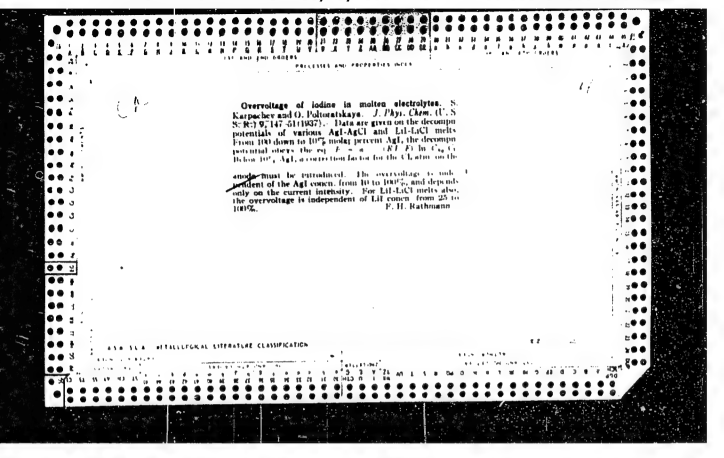


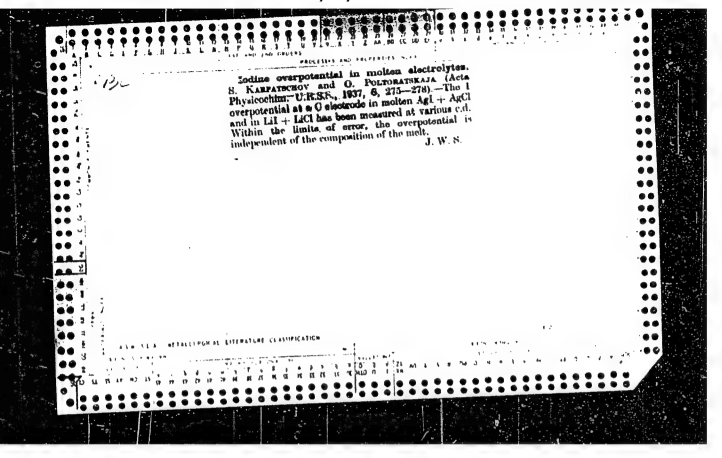


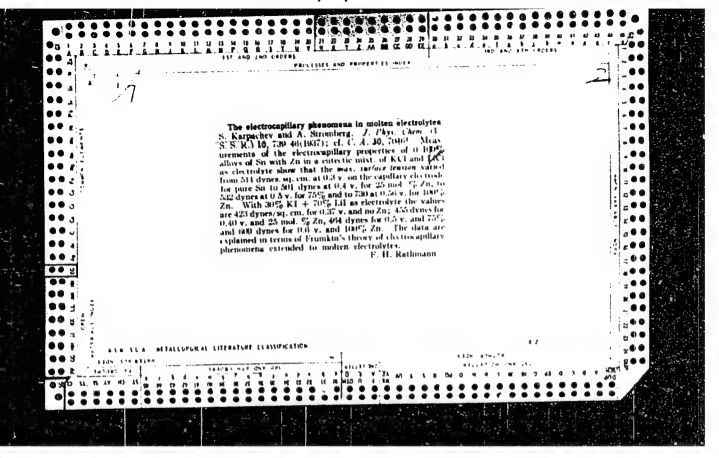


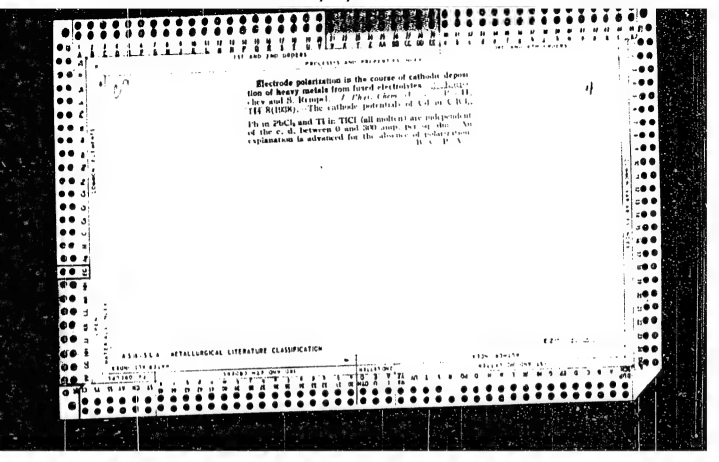


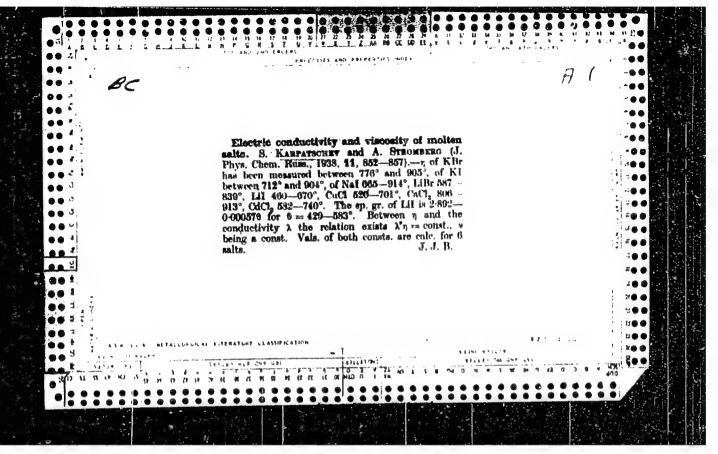








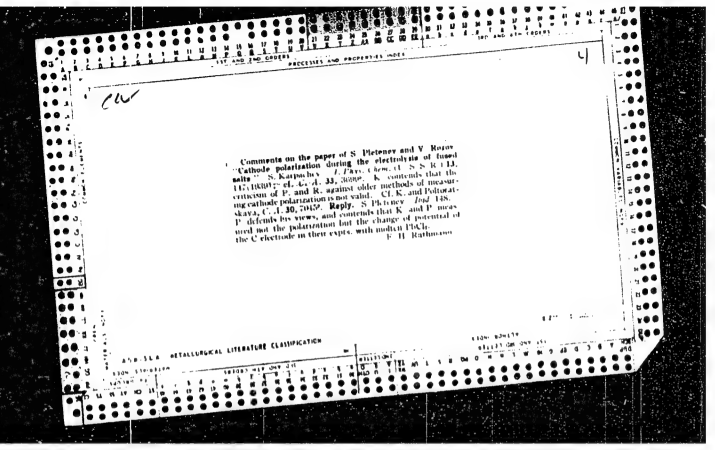


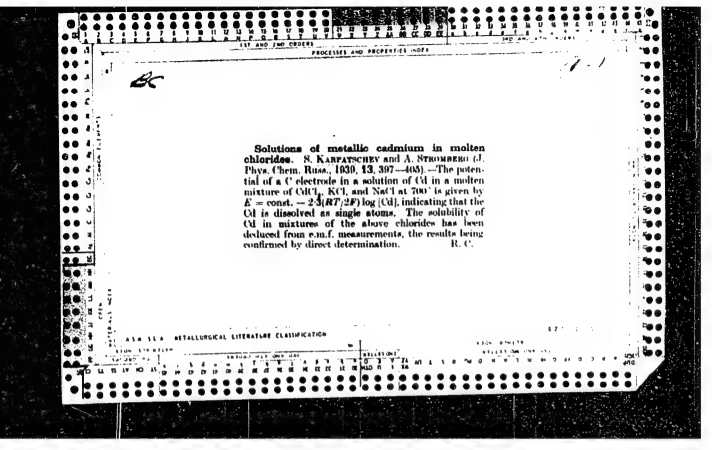


### KARPACHEV

"On the Work of S. Pletenev and V. Rozov, "Cathode Polarization during the Electrolysis of Molten Substances"; Zhur. Fiz. Khim., 12, No. 1, 1939.

Report U-1613, 3 Jan. 1952

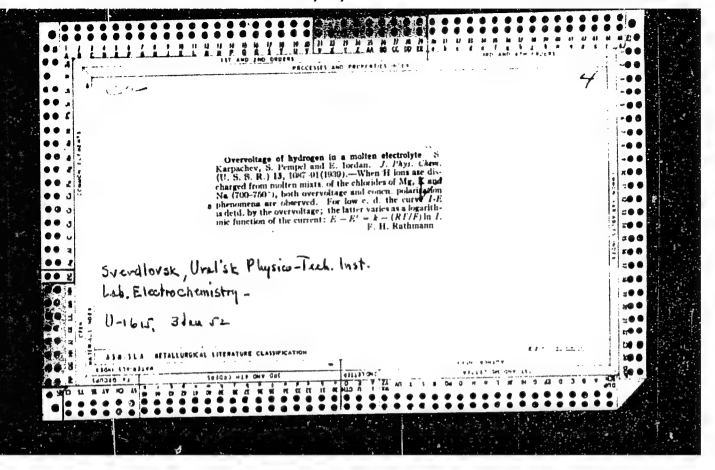


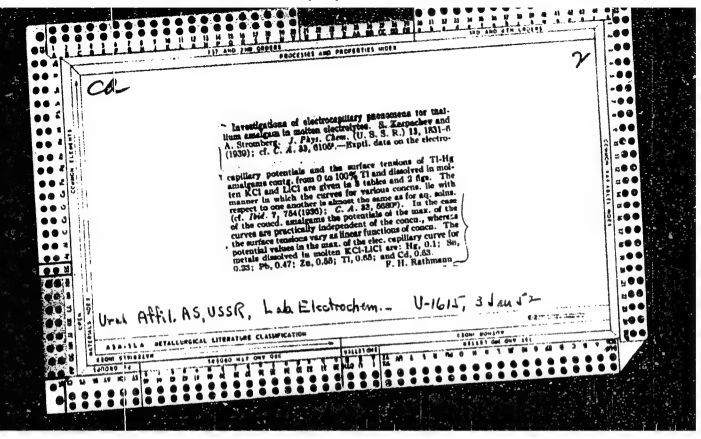


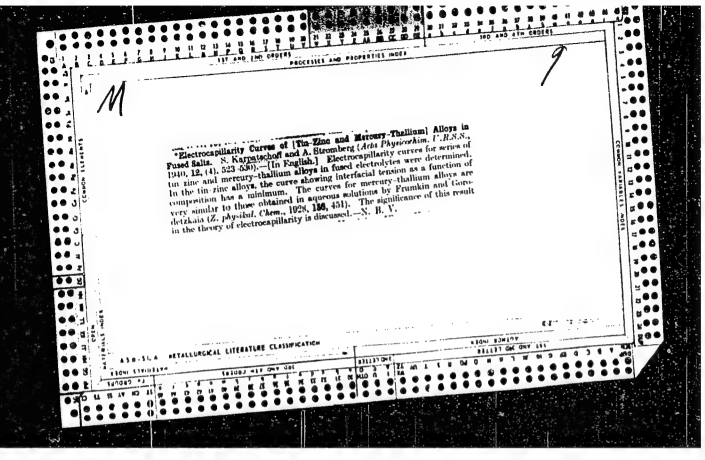
KARTACHEV, S., STROMBERG, A.

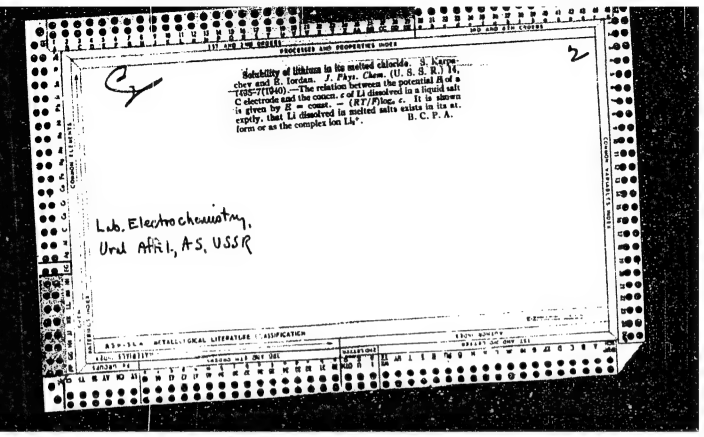
"On Solutions of Metallic Cadmium in Molten Chlorides"; Zhur. Fiz. Khim; 13, No. 3, 1939; Ural Physico-Tech. Insti., Lab. of Electrochem., Sverdlovsk; Rcd 28 Jul. 1938.

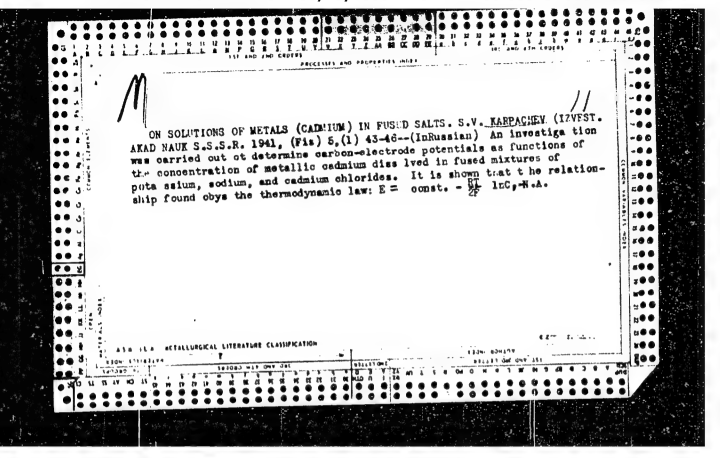
Report U-1.613, 3 Jan. 1952

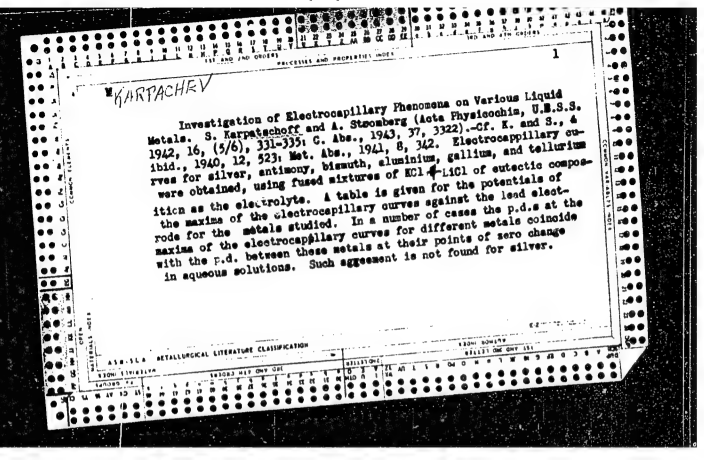


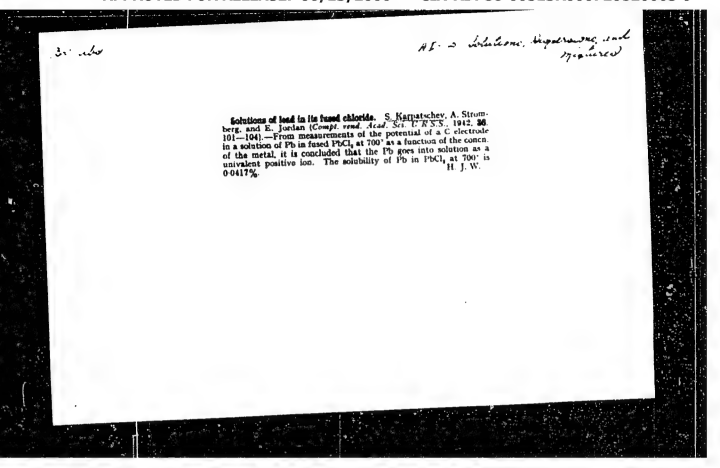










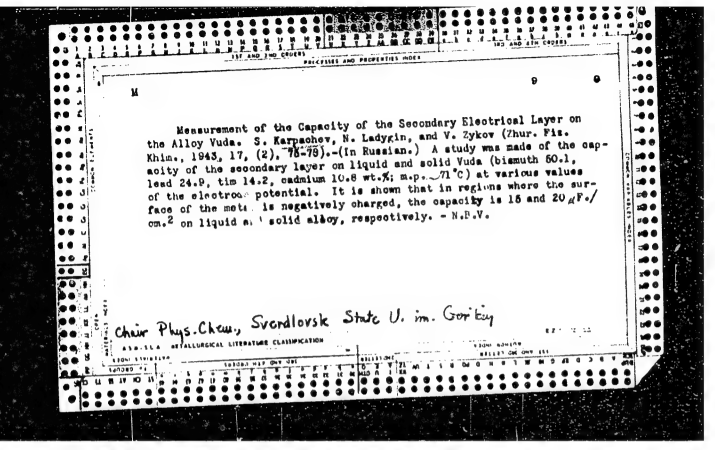


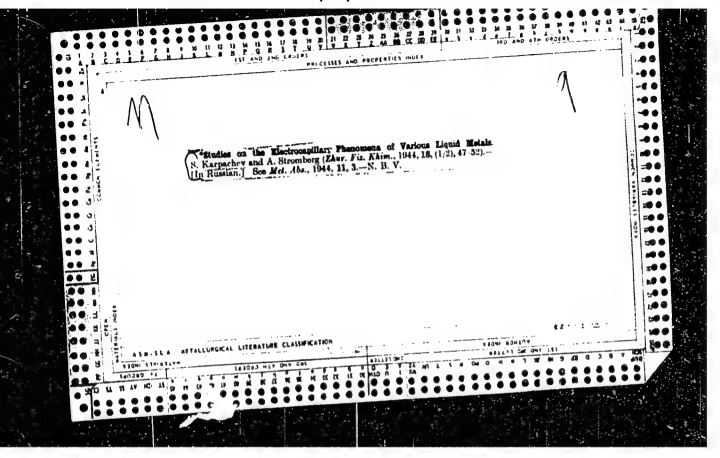
KARPACHEV, S., STROMBERG, A.

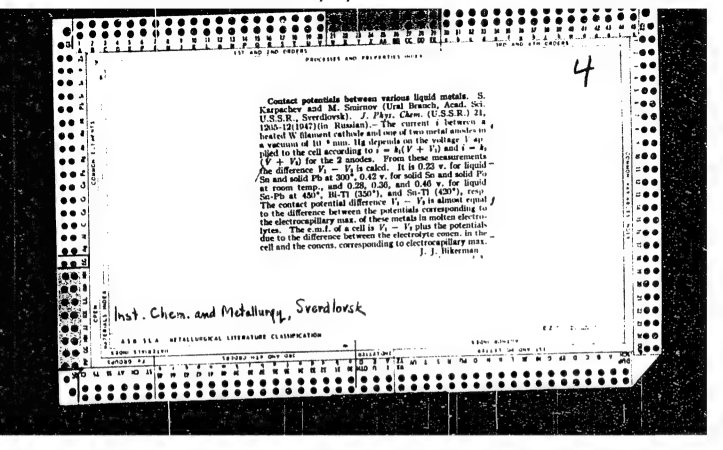
Mbr. Lab of Electrochemistry, Ural Affiliate of the Acad. of Sci., USSR; (-1941-)

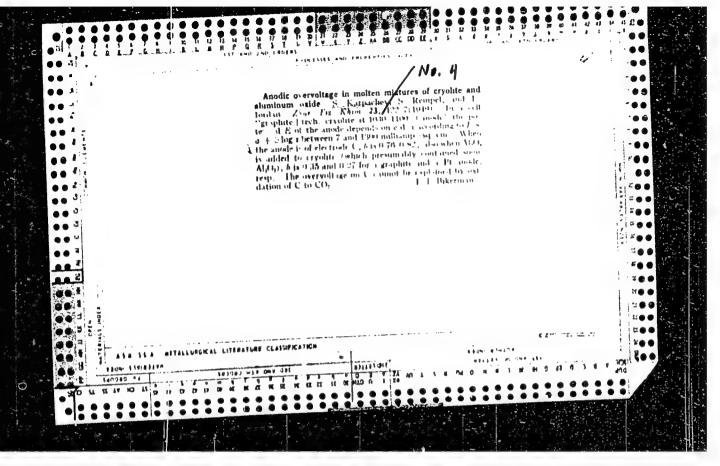
"The Contact Difference of Potentials Between Mercury and an Amalgam of Thallium" Zhur. Fiz. Khim. Vol 17, No. 1, 1943

BR-52059019



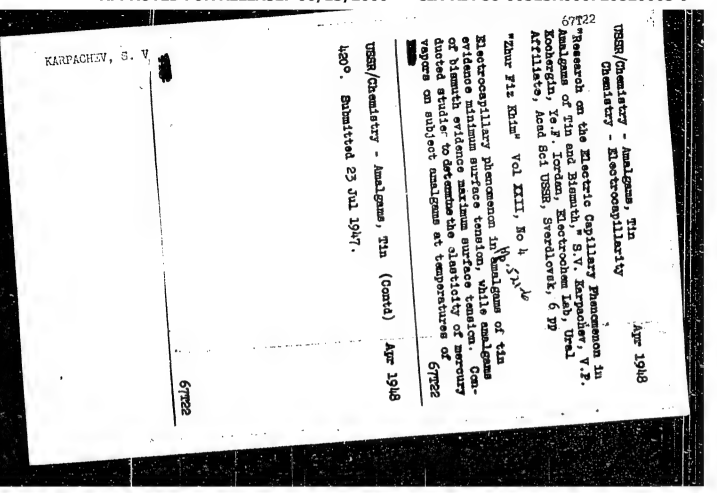






"APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000720820003-9



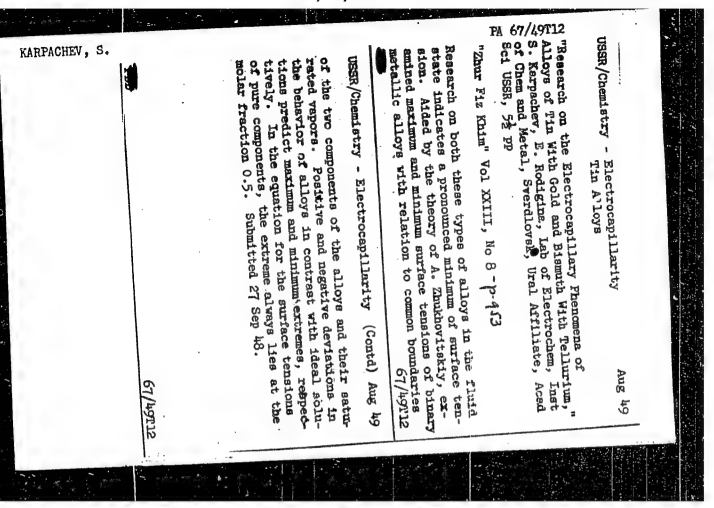
MARPOHIN, S.

Karpachev, . and Poliguev, S., Determination of the transfer numbers of cation and mion in rure melted calt. P. 942

The transfer numbers of the cation and anion in melted leaf chloride are measured with and without a porous diaphragm. Values for the transfer numbers of the cations and anion were obtained in an interval of 0.1-0.2 and 0.9-0.8 respectively; that is, the mobility of the cation in melted lead chloride is much smaller than that of the anion.

Leb. of Electrochemistry The Ural Branch of Acad. of Sci., USSR 5 pt. 3, 1948

30: Journal of Physical Chemistry (USSR) 23, No. 9 (1949)

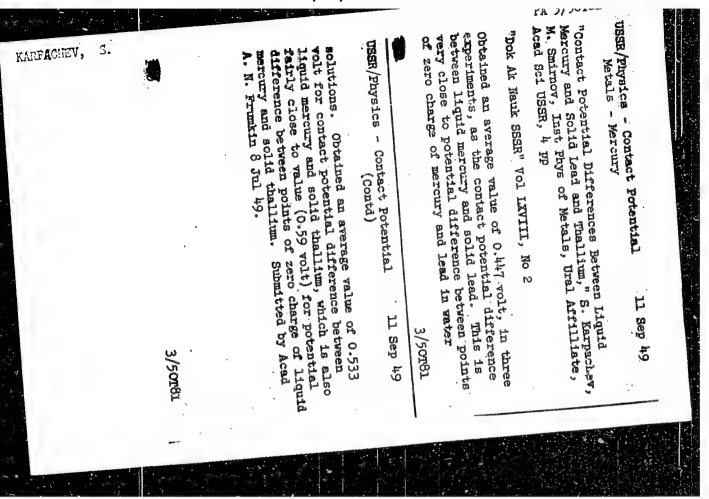


APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000720820003-9"

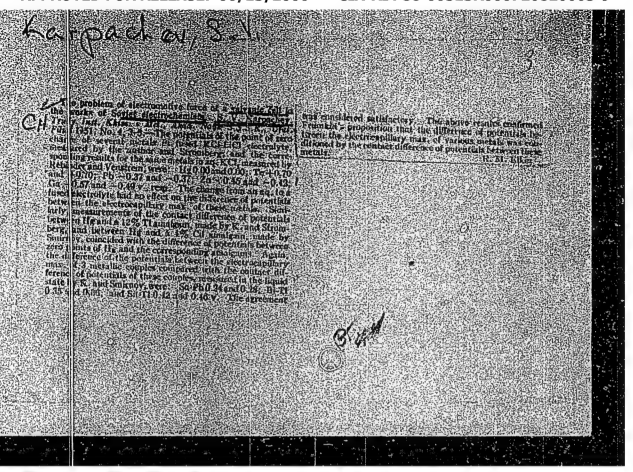
"APPROVED FOR RELEASE: 06/13/2000

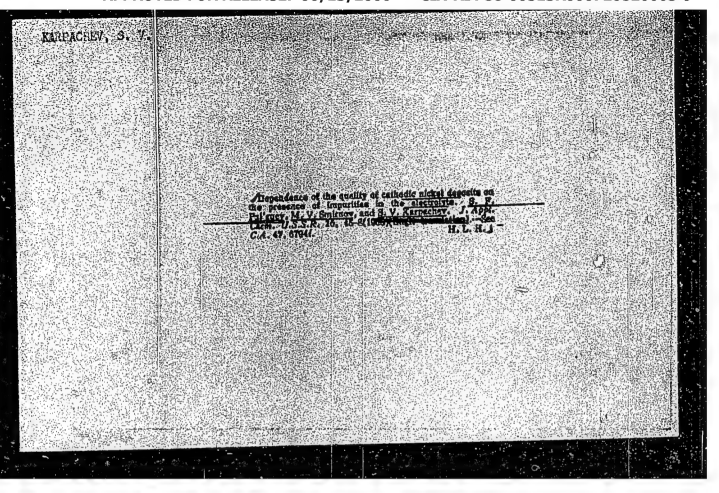
CIA-RDP86-00513R000720820003-9



APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000720820003-9"





### "APPROVED FOR RELEASE: 06/13/2000

### CIA-RDP86-00513R000720820003-9

USSR/Chemistry - Electrochemistry Maisling

KARPACHEV, S. V.

Jan 33

"te Boloti no ip letreem the quality of Nickel Cathole Beposito : 1 to Presence of I to white it to Electrolyte," S. 1. Pal' myev, F. W. Jahrnot, and S. W. Karpacher, Instead the listing and Petallury, Cont L. Miliate, Acad Sci USS

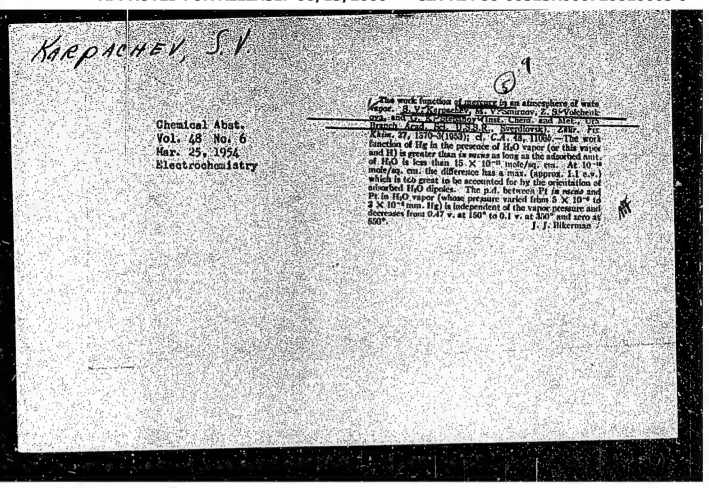
Thur Prit Khir, Vol 2', No 1, pp 50-51

Demonstrated the strong influence of the presence of one can be traces of moisture in the electrolyte on the nickel cathole isposit for intellectrolysis of rolten chloride. Passing dry hydrogen chloride through the rolt in electrolyte freed to that the first the above inpurities and improves the guilty of the stallic cathole jessit.

(CA 47 no: 14:6794 '53)

ŧ

2.7135



5(4) 5.4600

sov/20-129-6-39/69

AUTHORS:

Pal'guyev, S. F., Karpachev, S. V., Yushina, L. D.

TITLE:

An Electrochemical Chain With a Solid Electrolyte

PERIODICAL:

Doklady Akademii nauk SSSR, 1959, Vol 129, Nr 6, pp 1339-1342

(USSR)

ABSTRACT:

The authors investigated the temperature dependence of the electromotive force of the chain Ag/AgBr solid/Br,C in the region 25-421°. The electrochemical cell consisted of pressed silver bromide, to the one end of which a silver wire and to the other a carbon rod is attached. The entire cell was in a bromine atmosphere. Figure 1 gives the measured emf of the cell in dependence on temperature. The values between 110 and 421° lie on a straight line, for which an empirical equation is set up. Extrapolation to the melting point 434° of the AgBr gives an emf of 0.8033 v, which is in good agreement with the values given in reference 5 for liquid AgBr (0.8056 v), confirms the correctness of measurements, and points out that in solid AgBr electric conductivity near melting point has ionic char-

acter. The occurrence of electron conductivity in a bromine atmosphere is proven by the fact that the ion transfer number

Card 1/2

68172

An Electrochemical Chain With a Solid Electrolyte SOV/20-129-6-39/69

for Ag at 20° was between 0.03 and 0.05, and at 93° between 0.06 and 0.11, whereas that for Br was zero. On the basis of the thermodynamics of irreversible processes the concentration gradient of Br dissolved in solid AgBr and, thus, the gradient of the holes was calculated and the transfer number of the holes was found to be zero for the temperature interval of from 100 to 434°. The authors thank Academician A. N. Frumkin for the evaluation of their work. There are 1 figure and 8 references, 1 of which is Soviet.

ASSOCIATION: Institut elektrokhimii Ural'skogo filiala Akademii nauk SSSR (Institute of Electrochemistry of the Ural Branch of the Academy of Sciences, USSR)

PRESENTED: July 16, 1959, by A. N. Frumkin, Academician

SUBMITTED: July 15, 1959

Card 2/2

S/631/60/000/001/009/014 B110/B102

AUTHORS:

Karpachev, S. V., Vorob'yev, G. V.

TITLE:

Electrical conductivity of molten alkali-metal carbonates.

I. The system Na<sub>2</sub>CO<sub>3</sub>-K<sub>2</sub>CO<sub>3</sub>

SOURCE:

Elektrokhimiya rasplavlennykh solevykh i tverdykh elektrolitov,

no. 1, 1960, 61 - 64

TEXT: The authors studied the electrical conductivity of molten carbonates of the system Na<sub>2</sub>CO<sub>3</sub>-K<sub>2</sub>CO<sub>3</sub> between 700 and 1000°C. The end of an alundum tube (inner diameter: ~0.4 cm; wall thickness: 0.15 - 0.20 cm) was provided with cylindrical Pt electrodes with a surface area of ~0.5 cm². 0.5-mm Pt wires were used for current supply. At an electrode spacing of 4.0 - 5.0 cm, a 40 - 50 cm² cell was obtained, which was set by screws to an immersion depth of the upper electrode of 0.5 cm. The measuring arrangement consisted of an a-c bridge and a cathode-ray oscilloscope which was used as zero indicator. The conductivity was measured between 1000 - 5000 cps. The results were extrapolated to infinite frequency; Card 1/3

Electrical conductivity of ...

S/631/6**0**/000/001/009/014 B110/B102

 $R=g(1/\sqrt{f})$  (R= resistance, f= a-c frequency) is linear. The alundum tube remained unchanged during 8-hr melting of  $Na_2CO_3$  and  $K_2CO_3$  at  $900^{\circ}C$  in a  $CO_2$  atmosphere. The crucible with the alundum tube containing the melt was put into a metal block and then heated. The temperature was measured with a Pt-PtRh thermocouple. The temperature was kept constant with an accuracy of  $1-2^{\circ}C$  (measuring error,  $\frac{1}{2}$  3%). The temperature dependences of the electrical conductivities are straight lines which are almost parallel, since the temperature coefficients of the conductivity of pure, molten  $Na_2CO_3$  and  $K_2CO_3$  are similar. The curve of conductivity versus composition first ascends slowly when  $Na_2CO_3$  is added (with 40 - 4% of  $Na_2CO_3$  o has changed only by 4% and then rapidly up to the value of pure  $Na_2CO_3$ . The dependence of equivalent conductivity on composition is more complex: on melting with 50 mole% of  $Na_2CO_3$  a distinct minimum is observed. This indicates that the interaction between  $Na_2CO_3$  and  $K_2CO_3$  in melts reduces the conductivity and causes a density maximum at 45 mole% Card 2/3

Electrical conductivity of ...

S/631/60/000/001/009/014 B110/B102

of K<sub>2</sub>CO<sub>3</sub>. There are 2 figures and 7 references: 1 Soviet and 6 non-Soviet. The three references to English-language publications read as follows: E. Gorin, H. L. Recht. Amer. Soc. Mech. Engrs, 1958, A-200; A. Reisman. J. Amer. Chem. Soc., 81, 807, 1959; E. R. Van Artsdalen, J. S. Yaffe. J. Phys. Chem., 59, 118, 1955.

Card 3/3

### "APPROVED FOR RELEASE: 06/13/2000 CIA

CIA-RDP86-00513R000720820003-9

Electromotive forces of electrochemical cells with solid electrolytes. Trudy Inst.elektrokhim.UFAN SSSR no.1:79-89
160. (Electromotive force)
(Electrolytes)

29413 5/081/61/000/017/013/166 B102/B138

24,2130 (1142,1160,1482)

AUTHORS:

Karpachev, S. V., Pal'guyev, S. F.

TITLE:

Electromotive forces of electrochemical circuits with solid

electrolytes. II. Coexistence of n-type and p-type

conductivity in the electrolyte

PERIODICAL:

Referativnyy zhurnal. Khimiya, no. 17, 1961, 67, abstract 176495 (Tr. In-ta elektrokhimii. Ural'skiy fil. AN SSSR,

no. 1, 1960, 91-96)

TEXT: An expression is derived for the emf of a galvanic cell  $A/A_{z_2} X_{z_1}/X$ , C, A (A - metal of valency  $z_1$ , X - metalloid of valency  $z_2$ , C - carbon electrode, Az2, Xz1 - salt or oxide) taking account of ionic,

n-type, and p-type conductivity. It is shown that if n-type and p-type conductivity are due to dissolution of the metal or the metalloid in the solid electrolyte the emf of the circuits under consideration may, under certain conditions, be approximately the same as its thermodynamic value. [Abstracter's note: Complete translation.]

Card 1/1

29414

S/081/61/000/017/014/166 B102/B138

24,2130 (1142,1160,1482)

AUTHORS:

Karpachev, S. V., Pal'guyev, S. F.

TITLE .

Electromotive forces of circuits with solid electrolytes

possessing electron besides ion conductivity

PERIODICAL:

Referativnyy zhurnal. Khimiya, no. 17, 1961, 67-68, abstract

176496 (Tr. In-ta elektrokhimii. Ural'skiy fil. AN SSSR,

no. 1, 1960, 97-100)

TEXT: The effect of n-type conductivity on emf is considered for the electrochemical circuit Br,C/AgBr(solid)/Ag as an example. It is shown that  $E=E_0(1-t)$ , where E is the total emf,  $E_0$  the emf without n-type

conductivity, and t the number of electron transitions into the electrolyte. The efficiency of this kind of cell is described by the ratio:

 $((1-t)/(1-t^{(Ag)}))$ 100, if the efficiency of a cell without n-type conductivity is taken as 100 %. When current is obtained from the element, the Br distribution along the electrolyte changes, and the efficiency increases. [Abstracter's note: Complete translation.]

Card 1/1

KARPACHEV, S.V.; PAL'GUYEV, S.F.

Functioning of a primary cell with a solid electrolyte having both ionic and electron conductivity. Trudy Inst. elektrokhim.UFAN SSSR no.1:101-103 '60. (MIRA 15:2) (Electric conductivity)

PAL'GUYEV, S.F.; KARPACHEV, S.V.; YUSHINA, L.D.

Problem of measuring the decomposition voltage of solid electrolyte solutions. Trudy Inst.elektrokhim.UFAN SSSR no.1:105-110 '60. (MIRA 15:2)

(Solutions, Solid) (Electromotive force)

S/076/60/034/05/22/038 B010/B002

AUTHORS:

Karpachev, S. V., Karasik, E. M.

TITLE:

On the Heats of Fusion of Some Inorganic Salts

PERIODICAL:

Zhurnal fizicheskoy khimii, 1960, Vol. 34, No. 5,

pp. 1072-1076

TEXT: Proceeding from the concept of a salt melt as a crystal with a large number of lattice defects, the authors calculated the latent heat of fusion of some alkali halides (Table, results) by using Oriani's method of calculation (Ref. 4). The total change in entropy during the fusion process was represented as the sum from the change in entropy, caused by an increase in volume, and the change in entropy, caused by the increase in the degree of disorder on the fusion of the salt. The values of the latent heats of fusion calculated from this change in the entire entropy in the fusion process were compared with the corresponding experimental values taken from the manual by E. V. Britske, A. S. Kapustinskiy, and others (Ref. 11), and a satisfactory agreement was found (Table). Frenkel' is mentioned. There are 1 table and 12 references: 2 Soviet, 1 German, 6 American, 2 English, and 1 Japanese.

 $\sqrt{B}$ 

On the Heats of Fusion of Some Inorganic

S/076/60/034/05/22/038 B010/B002

Salts

ASSOCIATION: Uraliskiy gosudarstvennyy universitet im. A. M. Gorikogo

Sverdlovsk

(Ural State University imeni A. M. Gor'kiy, Sverdlevsk)

SUBMITTED:

July 19, 1958

Card 2/2

S/020/60/133/02/41/068 B004/B064

AUTHORS:

Karpachev, S. V., Pal'guyev, S. F.

TITLE:

On the Operation of a Voltaic Cell With a Solid

Electrolyte Having a Mixed Conductivity

PERIODICAL:

Doklady Akademii nauk SSSR, 1960, Vol. 133, No. 2,

pp. 396 - 398

TEXT: The authors investigated the effect upon the power of a voltaic cell of the electron conductivity of a solid electrolyte existing along with the ionic conductivity. The calculations were made from a sample cell with silver bromide as a solid electrolyte: Ag AgBr Br, C. The gaseous bromine should dissolve in AgBr and thus produce an electron conductivity of the p-type. On the condition that the bromine diffuses as far as the silver electrode and its concentration in the electrolyte is determined by the diffusion process only, the equations for the emf, the electron and hole transfer, the resistance of the electrolyte, and for the efficiency W without electron conductivity and for W with electron conductivity were derived (11). Equation (11) shows that in

Card 1/2

APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000720820003-9"

24.7700 1143, 1138, 1395

s/020/60/134/005/020/023 B004/B064

AUTHORS:

Pal'guyev, S. F., Karpachev, S. V., Neuymin, A. D.,

and Volchenkova, Z. S.

TITLE:

Transition From Electron to Ion Conductivity as a Function

of the Composition of Solid Solutions of Oxides

PERIODICAL:

Doklady Akademii nauk SSSR, 1960, Vol. 134, No. 5,

pp. 1138-1141

TEXT: The authors wanted to study the influence of calcium oxide upon the electrical conductivity of solid solutions of cerium and zirconium cxides. Since the solid solution 0.75CeO<sub>2</sub> °0.25ZrO<sub>2</sub> has the highest con-

ductivity (Ref. !), it was used as initial substance CaO was added in varying amounts; addition of CaO of up to 40 mole% resulted in the formation of solid solutions. The preparation of the samples was already described in Ref. 1. The relative electron and ion conductivities were determined by the solid electrolyte emf method at temperatures ranging from 500° to 1000°C, and herefrom the activation energy was computed. It was found that ion conductivity increases with an increase in tempera-

Card 1/4

Transition From Electron to Ion Conductivity S/020/60/134/005/020/023 as a Function of the Composition of Solid B004/B064 Solutions of Oxides

Card 2/4

Transition From Electron to Ion Conductivity S/020/60/134/005/020/023 as a Function of the Composition of Solid B004/B064

transference numbers of electrons and holes, respectively,  $E_0$  the thermodynamic value of the emf;  $E_0 = (RT/4F) \cdot \ln(P_2/P_1)$ . Table 1 lists the experimental data. Fig. 2 shows conductivity and  $\Delta E$  at  $1000^{\circ}C$  as a function of the CaO content. With rising CaO content in the system  $CeO_2 - ZrO_2 - CaO$ , the authors found a steady transition from electron to ion conductivity. This effect was not observed in the systems  $CeO_2 - CaO$  and  $ZrO_2 - CaO$ . The authors give the following explanation: As a result of partial reduction of  $Ce^{4+}$  to  $Ce^{3+}$ , first an intense electron conductivity occurs in the system  $CeO_2 - ZrO_2$ . Increasing addition of CaO inhibits this reduction more and more, and the ion conductivity caused by oxygen ions takes the place of electron conductivity. A decrease of conductivity in samples containing over 40 mole% CaO is attributed to the accumulation of free CaO not converted into a solid solution. There are 1 figure, 2 tables, and 9 references: 5 Soviet, 3 US, and 1 German.

Card 3/4

5.4700

S/631/61/000/002/013/01+ 1003/1203

**AUTHORS** 

Filyayev, A. T., Pal'guyev, S. F., and Karpachev, S. V.

TITLE

Investigation of electrode polarization in solid electrolytes

SOURCE

Akademiya nauk SSSR. Ural'skiy fiilial. Institut elektrokhimii Frudy, no 2, 1961-

Elektrokhimiya rasplavlennykh solevykh i tverdykh elektrolitov 199-209

TEXT. There are indications in the literature that the conductivity at elevated temperatures in a solid electrolyte consisting of 15 mole % CaO and 85 mole % of ZrO2 is almost entirely ionic and supposedly due to the movement of oxygen ions. The polarization of the CO + CO<sub>2</sub> (Pt) |solid electrolyte 0 15 CaO 0 85ZtO<sub>2</sub>| O2(Pt) cell was measured by a device containing a potentiometer or an oscilloscope. The oxygen electrode was used as the reference electrode, because it is not readily polarized. The relationship between the anodic polarization of the CO-CO2 electrode and the polarizing current density is expressed by the equation  $\mu = q + b \cdot \log i$ , where  $t \approx 2.3 \cdot RT/F$  for both anodic and cathodic polarization. Oscillographic measure ments show that, after the current is cut off, the polarization decreases so slowly that it can be correctly measured by means of an oscillograph. This is probable due to changes in concentrations caused by in hibition of diffusion or of chemical processes. There are 9 figures and 4 tables

Card 1/1

8/076/61/035/002/007/015 B124/B201

26.2520 AUTHORS:

Yushina, L. D., Paleguyev, S. F., and Karpachev, S. V.

TITLES

Study of electrochemical cells with solid electrolytes. I. Temperature dependence of the electromotive force of the cell  $Ag \mid AgBr(e) \mid Br_{n,0} \mid 0$ 

PERIODICAL: Zhurnal fizicheskoy khimii, v. 35, no. 2 1961, 342 - 349

TEXT: In the course of the work under consideration the authors measured the emf of the cell mentioned in the title in the range from room tempe-

rature to 421°C. The best results were obtained with a U-shaped glass tube; into whose one bend, which served as the tell proper, gaseous brownine was introduced and drawn off on the electrolyte surface. The contact with the bromine electrode in the tube was brought about by a soldered platinum wire. The bromine electrode used was a roulet of spectrally pure carbon and the electrolyte a sylinder pressed from AgBr. The emf of the cell was measured with a high-resistance potentiometer of the type NNTS-1 (PPTV-1), equipped with an Nt-91/A (M.91/A) galvanometer as a zero

Card 1/5

APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000720820003-9"

\$/U15/E /035/002/007/015 B 24/B201

Study of electrochemical cells ...

instrument. The measuring accuracy was  $\pm$  0. m: at temperatures over 110°C, and about  $\pm$  1.0 mm at lower temperatures. Measurement results at different temperatures are illustrated in Fig. 1. In the temperature range from 110 to 421°C the points established experimentally were upon a straight line, whose equation was derived empirically with the aid of the least squares method:  $E = (1.4518 - 0.493 \cdot 0^{-3}T) \pm 0.0017 \cdot v$  (1); at temperatures lower than  $110^{\circ}C$  the points established experimentally were not on a straight line. According to Ref. 1 (E. J. Salstrom, J. H. Hildebrand, J. Amer. Chem. Soc. 52, 5650. 1930%, the emf of the cell with liquid AgBr amounts to 0.8056 v at 434°C while the one according to Eq. (1) is 0.8033 v. By thermodynamic calculations made on the basis of data by A. F. Kapustinskiy (Ref. 8). Termicheskiye konstanty neorganicheskikh veshchestv" (thermal constants of inorganic substantes) M. - L.: 1949) 0.9904 v was found for the emf at 25°C, while 0049 v was obtained from Eq. (1). These data prove that there exists practically no

Card 2/5

S/075/61/035/002/007/015 B124/B201

Study of electrochemical cells ...

electron conductivity in AgBr in the temperature range from 420 to 100°C. To find out whether the electrical conductivity of solid AgBr was of an electronic nature, the authors measured the transfer numbers of the ions in solid AgBr in bromine (vapor pressure = 168 mm Hg) at room temperature and 93°C, the above described device being used for this purpose also. The electrolyte plates used were pressed by a special equipment between the carbon electrodes. The electrolysis current at 20°C was 0.5 to 1.2 ma and at 93°C it was 1.0 to 5.0 ma. The dissolution of bromine in solid AgBr causes an electron conductivity to appear. The electron conductivity arising in solid AgBr in connection with the bromine dissolution is caused by the motion of positively charged holes in the electric field. Ag- and Br ions and positively charged holes were found in solid AgBr. The mean value of the hole-transfer number is calculated from equation

 $t_o = (\int_{Br}^{Ag} t_o \ d\mu_{Br})/(\mu_{Br}^{Ag} - \mu_{Br}^{Br}) \ (11) \ (\text{where} \ \mu_{Br} \ \text{denotes the potential of}$  atomic bromine in bromine vapors around the Br=electrode,  $\mu_{Br}^{Ag}$  and  $\mu_{Br}^{Br}$  the chemical potential of atomic bromine, dissolved in the crystal around the

Card 3/5

S/076/61/035/002/007/015 Study of electrochemical cells ... B124/B201

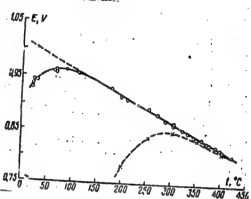
Ag- and Br-electrode, respectively; to is proportional to the concentration of atomic bromine dissolved in AgBr: to - kcBr. The relation  $t_o = \left[k(c_{Br}^{Br} - c_{Br}^{Ag})\right] / \left[ln(c_{Br}^{Br}/c_{Br}^{Ag})\right]. \text{ The relation } t_o^{Br}/c_{Br}^{Ag} / ln(c_{Br}^{Br}/c_{Br}^{Ag})$ (12) is derived, where to is the hole-transfer number in AgBr saturated with bromine at a bromine vapor pressure of 170 mm Hg, and cBr is the concentration of dissolved bromine in the immediate neighborhood of the Br elec-It may be stated in this connection that if the solid salt saturated with a metalloid practically exhibits only an electron conductivity, this will not become manifest at very low concentrations of dissolved solid metalloid salt near the metal electrode when measuring the emf; the same holds for dissolution in a solid metal salt. There are 1 figure and 10 references: 2 Soviet-bloc and 8 non-Sovietbloc. A reference to English language publications reads as follows: D. L. Hildebrand, W. R. Kramer, R. A. Mac Donald, D. R. Still, J. Amer. Chem. Soc., 80, 4129, 1958. ASSOCIATION: Institut elektrokhimii, Sverdlovsk (Institute of Electrochemistry, Sverdlovsk) Card 4/5

Study of electrochemical cells .

SUBMITTED: May 30, 1959
Legend to Fig. 1: Temperature dependence of the AsiAgBr(sol) iBr C college

AglAgBr(sol) Br., C cell;
1) data obtained from the present work; 2) data calculated on the basis of paper Ref. 3: C. Wagner, Z. phys. Chem., 32, 447, 1936.

09573 8/076/61/035/002/007/015 B124/B201



Температурная зависямость цени Ag [AgBr(тв) Brs, С: — данные настоящей работы; 2 — данные, рассчитавные на основании работы [3]

Card 5/5

Card 1/4

3070**6** \$/020/61/141/002/020/027 B101/B147

Polarization of some gas ...

which could become constant (measurement with a Pt-PtRh thermocouple). The resistance of the electrolyte was measured with an a-c bridge (3000 cps). At the experimental temperatures 900, 1000, and 1100°C, it was 23.90, 7.93, and 3.80 ohms, respectively. A linear relationship was found between current density and the voltage measured at the electrodes. The resistances of the electrolyte calculated from the slope of the straight line showed only slight divergences from the values measured directly. Accordingly, no overvoltage occurred at the 02 electrode. Then, a Pt,02 electrode was exchanged for a Pt, CO + CO2 electrode (cell II) which produced a noticeable emf, so that no external current source was necessary. Amperage was changed by means of a resistance box. The anodic polarization of the CO electrode around which a flow of 66 vol; of CO + 34 vol% of CO2 streamed, was then measured. Pure oxygen streamed around the  $O_2$  electrode. The anodic overvoltage was determined from  $\eta = E_0 - Ir - IR$  (1).  $\eta = absolute value of anodic overvoltage;$  $E_0 = equilibrium value of emf; I = amperage; r = resistance of electrolyte$  $I\bar{R}$  = potential drop in the external circuit. For all of the three temperatures, the linear function  $\eta = a + b \log i$  (2) was found (1 ... pressed in a/cm2). Since two electrons participate in the anodic p. Card 2/4

Polarization of some gas ...

\$/020/61/141/002/020/027 B101/B147

of the CO electrode,  $b = 2.3RT/2F\alpha$  is written. For a, b,  $\alpha$  the following values are given:

	900°c	1000°C	1100°C
Ъ	0.258	0.250	0.260
α	0.45	0.51	0.52
a	1.068	0.953	0.866

At low amperages, the measurement values deviated considerably from linearity. This was ascribed to the polarization countercurrent. The following equations are written:

 $i = K \exp(\eta F/RT)$  (3);  $i = K \exp(-\eta F/RT)$  (4); i = i + i (5). At high overvoltage,  $i \approx i$ , at low overvoltage Eq. (5) is valid. i is the current measured. By using Eq. (4), i was calculated for the measuring points deviating from linearity at low overvoltage, and, by taking i into account, complete linearity could be achieved also for these measuring points. At the CO electrode, a considerable overvoltage ( $\eta = 0.6 \text{ v}$ ) occurred. Thus, the electrodic reaction is inhibited in spite of the high temperature. Further studies are necessary to explain this effect. There

Card 3/4

ED FOR RELEASE: 06/13/2000

KARPACHEV, S.V.; FILYAYEV, A.T.; PAL'GUYEV, S.F.

inodic polarization in solid electrolytes. Trudy Inst. elektro-khim. UFAN SSSR no.3:161-164 '62. (MIRA 16:6)

(Electrolytes) (Polarization (Electricity))

FILYAYEV, A.T.; KARPACHEV, S.V.; PAL'GUYEV, S.F.

Capacitance of a double electric layer in solid electrolytes.

Dokl. AN SSSR 149 no.4:909-910 Ap '63. (MIRA 16:3)

l. Institut elektrokhimii Ural'skogo filiala AN SSSR. Predstavleno akademikom A.N.Frumkinym.

(Fused salts—Electric properties)

KARPACHEV, S. V.; FILYAYEV, A. T.; PALGUYEV, S. F.

\*Polarization of Oxygen Platinum Electrodes in a 0.85 ZrO2 0.15 CaO Solid Electrolyte.\*

Report presented at the 14th meeting CITCE, Intl. Comm. of Electrochemical Thermodynamics and Kinetics, Moscow, 19-25 Aug 63.

Ural Institute of Electrochemistry, Academy of Sciences of U.S.S.R.

48970=65 ENT(m)/EWG(m)/T RWH CESSION NR: AP5007751	\$/0364/65/001/001/0090/0093 /3
THOR: Perfil'yev, M. V.; Fal'guyev, S.	E.; Karpachev, S. V.
TLE: Impedance of the electrode-electro	lyte interface and the double layer capa.
OURCE: Flektrokhimiya, y. 1, no. 1, 196	i, 90-93
PIC-TAGS: electrode, electrolyte, plat	inum, impedance
f the type Pt. On solid electrolyte On ange. The electrolyte was a solid solute the form of a cylinder with pointed come electrode was platinum applied as power auxiliary electrode was applied on the other measurements the electrode was treeted.	de processes in solid electrolytes. Certs Pt were used in the 700-800°C temperature ion of 0:15 Ca0.0.85CeO2. This was made nical depressions at the ends. The operat- der on the surface of the central cylinder. external surface of the cylinder. Prior

L 48970-65 CCESSION NR: AP5007751 cycles with a voltage amplitude in the cell of 20 millivolts. The resistance of the electrolyte was measured with a cathode oscillograph. The resistance was 35-40 R and the accuracy of measurement was on the order of 1-2 R. All measurements were made at the equilibrium potential of the oxygen electrode being investigated. The results of the measurements are shown in table 1 of the Enclosure where  $\Delta \phi$  is in millivolts and C is in  $\mu f/cm^2$ . Orig. art. has: 2 figures, 1 table, 4 equations. ASSOCIATION: Institut elektrokhimil Ural'skogo filiala Akademii nauk SSSR (Institute of Electrochemistry, Ural Branch, Academy of Sciences SSSR) 01 SUB CODE: GC, IC ENCL: SUBMITTED: 01Sep64 OTHER: 003 NO REF SOV: 008

## "APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000720820003-9

AGC NR: AP5025351	IJP(c) JD/WW/JG/WH SOURC	)/EPF(n)-2/EPA(w)-2/EWP(t)/ E CODE: UR/0131/65/000/010/	0040/0042
		V.; Neuymin, A. D.; Pal'guy	
- VIIIIII	projection is then belonded I have made to be seen	v., neuymin, n. D.; rei-guy	
		<b>44</b>	77
ORG: Institute of elektrokhimii	Electrochemistry, Urals I UFAN SSSR)	Branch, AN SSSR (Institut	W)
CITLE: Penetration	of oxygen in ceramics ha	ving a zirconium dioxide ba	59
SOURCE: Ogneupory,	no. 10, 1965, 40-42	31	15, 44
COPIC TAGS: ceramititani	c product, gas diffusion,	oxygen, carbon monoxide, a	rgon,
laving a composition the flow of oxygen - with argon and tital	n of 0.85+ZrO <sub>2</sub> 0.15 CaO, that diffused through the nium. A test was also car	it of oxygen diffusion throu in the temperature range of walls of the test tubes wa ried out to determine the p It was less than 4 • 10	600-900°C. s measured

# ACC NR: AP5025351 The ceramics tested proved to be practically impenetrable to carbon monoxide, and oxygen diffused directly through the ceramic body and not through the pores. This work was based on the experiments of Kingery W. D., Pappis J., Doty M. E., Hill D. C. Journ. Amer. Cer. Soc., 1959, v. 42, no. 8, p. 393. Orig. art. has: 3 figures and 1 table. SUB CODE: MT, GC SUBM DATE: 400/ NR REF SOV: OOI/ OTHER: OO2

OVCHINNIKOV, Yu.M.; KARPACHEV, S.V.; PAL'GUYEV, S.F.; ZHDANOVA, G.M.; NEUYMIN, A.D.

Kinetics of the reduction by carbon monoxide of solid solutions based on cerium dioxide. Elektrokhimiia 1 no.10:1196-1201 0 '65.

(MIRA 18:10)

1. Institut elektrokhimii Ural'skogo filiala AN SSSR.

VOROBITEV, G.V.; PALIGUYEV, S.F.; KURRYLEROV, S.V.

Viscosity of molten glasti metal carbonstes. Trudy Inco.
clartecokhim. UNAN SSSN no.5559-45 165. (MIRT 18.21)

OVCHINNIKOV, Yu.M.; KARPACHEV, S.V.; NEUYMIN, A.D.; PALIGUTEV, S.F.,

Oxygen permeability of ceramics on a 7rO<sub>2</sub> base. Ognoupory 50 no.10:40-43 165. (MIRA 18.10)

1. Institut elektrokhimii Uraliskogo fillala AN SSSR.

30219-66 ENT(m)/ETC(f)/T/ENP(t)/ETI IJP(c) DS/NW/JD/JG
ACC NRI AP6015014 (A) SOURCE CODE: UR/0364/66/002/005/0617/0615

AUTHOR: Karpachev, S. V.; Filyayev, A. T.

40 B

ORG: Institute of Electrochemistry, Ural Affiliate, Academy of Sciences SSSR, Sverdlovsk (Institut elektrokhimii Ural'skogo filiala Akademii nauk SSSR)

TITLE: Some specific features of the behavior of gas electrodes in contact with a solid electrolyte

SOURCE: Elektrokhimiya, v. 2, no. 5, 1966, 617-619

TOPIC TAGS: zirconium compound, calcium oxide, solid solution, electrolyte, platinum, electrode potential, cathode polarization

ABSTRACT: The vauthors measured the capacity of the electric double layer in the solid solution 0.852ZrO2·0.15CaO in order to refine the data obtained earlier and to use the method of capacity measurement for studying the behavior of gas electrodes in contact with a solid electrolyte. Platinum electrodes were employed and the capacity and resistance of the electric double layer were measured (at a frequency of 100 Kc) as functions of the electrode potential. Pretreatment of the electrode with anodic current decreased the electrode capacity. This is attributed to a decrease in the contact area between the electrolyte and the electrode, caused by the disrupting action of the oxygen escaping from the electrode, as indicated by a rise of the ohmic resistance and

UDC: 541.135.4

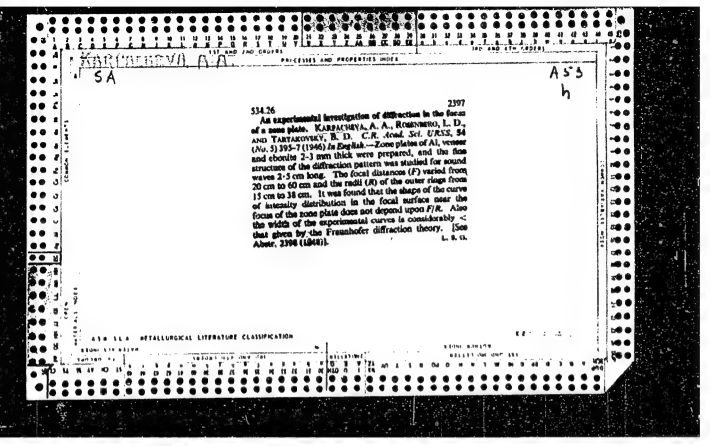
Card 1/2

· .
• :

#### "APPROVED FOR RELEASE: 06/13/2000

CIA-RDP86-00513R000720820003-9

38369-66 EWT(m)/T (A) ACC NR: AT6021374 SOURCE CODE: UR/2631/65/000/007/0169/017L AUTHOR: Filyayev, A. T.; Karpachev, S. V.; Pal'guyev, S. F. ORG: none TITLE: Study of the polarization of the oxygen electrode in a solid electrolyte SOURCE: AN SSSR. Ural'skiy filial. Institut elektrokhimii. Trudy, no. 7, 1965. Elektrokhimiya rasplaviennykh solevykh i tverdykh elektrolitov; termodinemika i kinetika elektrodnykh protsessov (Electrochemistry of fused salts and solid electrolytes; thermodynamics and kinetics of electrode processes), 169-174 TOPIC TAGS: electric polarization, oxygen, platimum, zirconium compound, calcium compound, electric impedance, electrode ABSTRACT: Continuing their investigation of electrode polarization in solid electrolytes, the authors studied the polarization of the platinum oxygen electrode as a function of current density in the solid solution 0.85Zr02.0.15CaO. The anodic polarization was determined from oscillograms of the electrode potential drop with time after the polarizing current was switched off. At relatively low polarizations, the current dependence of the polarization obeyed Tafel's equation,  $\Delta \phi = a + b \log i$ , where  $b = 1.3 \frac{2.3 \text{ RT}}{2.3 \text{ RT}}$ . It is shown that this value of the coefficient is compatible with the slow discharge theory. The impedance of the electrode - electrolyte inter-1/2



DAVYEDY, B.E.: MARHARYAN, B.Z.; KARPACHEVA, G.P.; KRENTSELI, B.A.; LAPITSKIY, G.A.; KHUTAREVA, G.V.

Disarrangement of complanarity and conjugation in crystalforming prlymers. Dokl. AN SSSR 160 no.3:650-653 Ja 165. (MIRA 18:3)

1. Institut neftekhimicheskog: sinteza im. A.V. Topchiyeva AN SSSR. Submitted July 14, 1964.

14.7000

39**1.7**4 5/056/62/043/002/002/053 B102/B104

AUTHORS:

Volkov, D. I., Pshenichkin, P. A., Karpacheva, G. P.

TITLE:

Temperature dependence of the magnetic susceptibility of

manganese-copper alloys

PERIODICAL:

Zhurnal eksperimental noy i teoreticheskoy fiziki, v. 43,

no. 2(8), 1962, 370 - 375

TEXT: The magnetic properties of Cu-Mn alloys show peculiarities, the causes of which have so far not been fully clarified. The authors studied (T) between 77 and 1250°K for Mn concentrations from 5.76 - 91.2 atts. Measurements were made by the weighing method. The alloys were fused in an evacuated h-f furnace. At low temperatures, all alloys show an antiferromagnetic transition (Neel point  $\theta_N$ );  $\theta_N$  lies the deeper, the lower the Mn content; from 50% Mn upward, it remains constant at about 150°K. The course of the curves  $\chi^{-1}(T)$  showing a minimum at  $\theta_N$  depends on the Mn content: up to 17%, they are linear between  $\theta_N$  and the melting point, Card 1/2

Temperature dependence of the ...

S/056/62/043/002/002/053 B102/8104

as from 22% between  $\theta_N$  and room temperature. They show a salient point at room temperature, and go on linearly up to the melting point. The inclination of these straight lines is the greater, the higher the line concentration. Resistivity maxima occur at these critical temperatures. Alloys containing 82.5 and 91.2 at/2 Mn show an abnormal behavior of  $\theta_N$  (2):  $\theta_N$  grows with the temperature. The experimental results can be explained by assuming that conduction electrons participate in the exchange interaction. There are 5 figures.

ASSOCIATION: Moskovskiy gosudarstvennyy universitet (Loscow State University)

SUBMITTED: January 8, 1962

Card 2/2

L 16034-65 ENT(m)/EPF(c)/EWP(1)/T Pc-4/Pr-4 AFWL/SSD/ASD(m)-3/AS(mp)-2/AFETP/ACCESSION NR: AP4045800 RAEM(a)/ESD(t)\$/0062/64/000/009/1697/1700

AUTHORS: Nasirov, F.M; Karpacheva; G.P.; Davy\*dov, B.E.; Krentsel!, B.A.

TITLE: Structure of the soluble complex organometallic catalyst for acetylene polymerization

SOURCE: AN SSSR. Izv. Seriya khimicheskaya, no. 9, 1964, 1697-1700

TOPIC TAGS: acetylene polymerization catalyst, complex organometallic catalyst, structure, chemical behavior, triethylaluminum, vanadium acetylacetonate, triethylaluminum vanadium acetylacetonate catalyst, tetravalent vanadium, divalent vanadium, magnetic susceptibility, EPR spectrum, magnetic moment, g-factor

ABSTRACT: The structure and the chemical nature of the active centers of the acetylene polymerization catalyst complex formed by reaction of triethylaluminum with vanadium acetylacetonate were examined. The catalyst, prepared by mixing VC10H1, 05 with a fourfold excess of A1(C2H5)3 in benzene at room temperature, appeared homogeneous. It was proposed that the formation of the active catalytic complex took place according to the reaction shown in the Cord 1/3

L 16034-65

ACCESSION NR: AP4045800

enclosure in which the tetravalent vanadium was reduced to the divalent. The magnetic susceptibility and the EPR spectra of the vanadium acetylacetonate and of the complex were examined. The magnetic moment for VC10H1.05, determined from the reverse molar magnetic susceptibility-temperature (120-300K range) relationship, was 1.67; for the complex, 3.83. Similar values for magnetic moment were calculated from g-factors obtained from EPR spectral date, confirming divalency of the vanadium in the complex. Orig. art. has: 3 figures and 2 equations.

ASSOCIATION: Institut neftekhimicheskogo sinteza im. A.V. Topchiyeva Akademii nauk SSSR (Institute of Petrochemical Synthesis Academy of Sciences SSSR)

SUBMITTED: 27Jan64

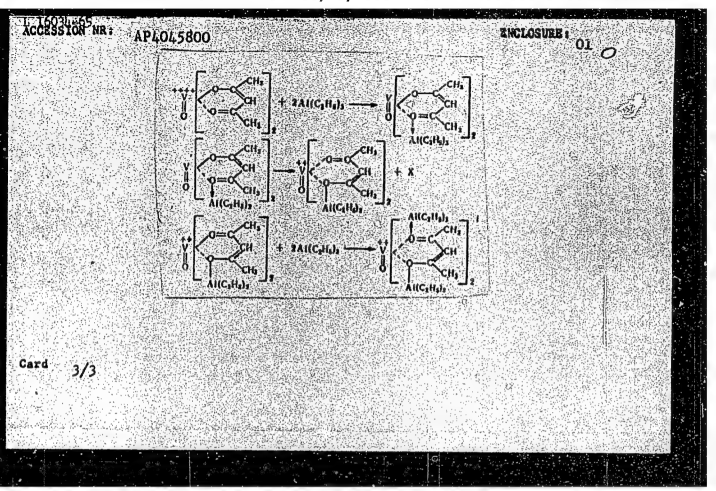
ENCL: Ol

SUB CODE: GC

NR REF SOV: 002

OTHER: 004

Card 2/3



L 54617-65 EPA(s)-2/ENI(d)/EPP(c)/ENP(j)/T, ENA(c) Pc-4/Pr-4/Pt-7 ACCESSION NR: AP5006423 \$/0062/65/000/001/0190/0192 AUTHOR: Karpacheva, G. P. TITLE: Investigation of systems of Schiff polybasic compounds and certain halides with charge transfer SOURCE: AN SSSR. Izvestiya. Seriya khimicheskaya, no. 1, 1965, 190-192 paramagnetic resonance, polymer, glyoxal, charge transfer, polycondensation ABSTRACT: Magnetic and electrical properties were studied in systems where the electron donor is the molecule of a polymer which belongs to the class of Schiff polybasic compounds; and the electron acceptors are bromine and iodine. The material used in this study was the product of polycondensation of glyoxal/with 2,6-diaminopyridine (with a molecular mass of 800. Complexes with donor-to-acceptor ratios of 1:0.4, 1:0.95; 1:4.5, 1:2, 1:3.3, 1:5 and 1:8.7 were studied. The intensity of the paramagnetic resonance signal, the resistivity, and conductance activation energy are each shown as a function of the bromine content in fig. 1 of the Enclosure. The resistivity is shown as a function of temperature Card 1/4

L 54617-65

ACCESSION NR: AP5006423

for each of the molar ratios in fig. 2 of the Enclosure. The identical nature of the variation in the three parameters (p, I, AB) and the coincidence of the extrema on the curves for the relationship between these parameters and the composition of the complex indicate that the nature of electrical conductivity and paramagnetism is the same and is associated with the formation of a complex with charge transfer. "In conclusion I consider it my duty to express my gratitude to D. A. Blyumenfel'd, B. E. Davydov and B. A. Krentsel' for their constant interest in the work." Orig. art. has: 2 figures.

ASSOCIATION: Institut neftekhimicheakogo sinteza im. A. V. Topchiyeva Akademii nauk SSSR (Institute of Petrochemical Synthesis, Academy of Sciences SSSR)

SUBMITTED: 03Jun64

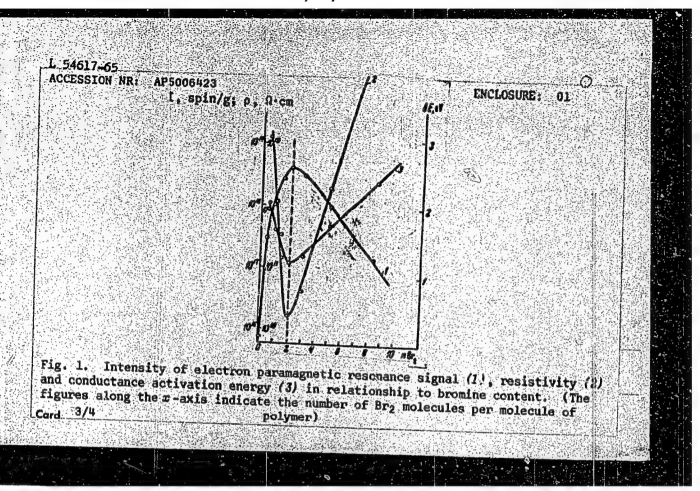
ENCL: 02

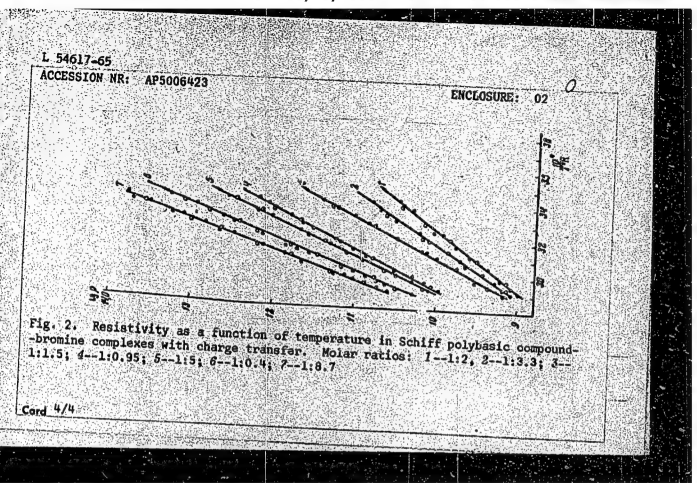
SUB CODE: GC, EM

NO REF SOV: 005

OTHER: 003

Card 2/4





L 29134-65 EPA(s)-2/EHT(m)/EPF(c)/ENP(j)/T Pc-4/Pt-10 RM

ACCESSION NR: AP5005899

8/0020/65/160/003/0650/0653

AUTHOR: Davydov, B. E.; Zekharyen, R. Z.; Karpscheve, C. P.; Krentsell, B. A.;

Lapitskiy, C. A.; Khutareva, C. V.

TITLE: Impairment of coplanarity and conjugation in crystallizing polymers

SOURCE: AM SSSR. Doklady, v. 160, no. 3, 1965, 650-653

TOPIC TAGS: crystallization, conjugation, conjugated polymer, organic semiconductor; semiconducting polymer, coplanarity

ABSTRACT: A study has been made to determine to what extent crystallization gives rise to conjugation disruption due to impairment of coplanarity in conjugated polymers in the solid phase, and how it affects their optical, paramagnetic, and semiconducting properties. These properties were compared for 32 polyazines and polymeric Schiff bases. It was found that the properties which are typical of conjugated polymers are exhibited to a greater extent by amorphous than by crystalline polymers. Thus, in color, in IR spectra, and in the absence of EPR, crystalline polyazines are similar to their analogs containing 0, 8, CH3, or OCH3 groups between conjugated segments in the backbone. A similar correlation, but less marked, was in evidence for the polymeric Schiff bases. This effect of crystallinity on con-

Cord 1/2

L 29134-65 ACCESSION NR: AP5005899 jugated-polymer properties was attributed to the impairment of coplemerity during crystallization. In thermal stability and activation energy for conduction, however the crystalline polymers were closer to the amorphous ones. The effect of crystallinity on semiconducting properties was interpreted as being determined in each individual case by changes in activation energy due to two competing processes occurring on crystallization: an increase in carrier mobility and a decrease in carrier concentration. Orig. art. has: I table. ASSOCIATION: Institut neftekhimicheskogo minteza imeni A. V. Topchiyeva Akademii nauk SSSR (Institute of Petrochemical Synthesis, Academy of Sciences, SSSR) BUB CODE: 55, OC SUBMITTED: 30Jun64 ENCLY 00 000 NO REF SOV ATD PRESS: 3197 OVHER : 000 Card 2/2

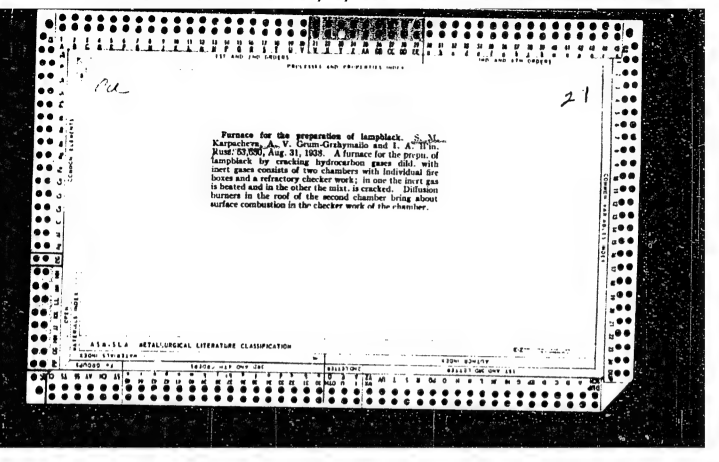
KARPACHEVA, S.M., doktor khimich. nauk; CHEMARIN, N.G., kand.tekhn.nauk;

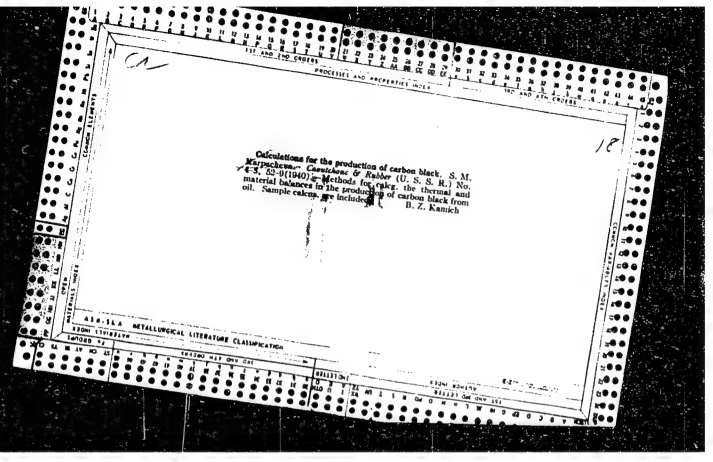
BYCHKOV, A.Ye., inzh.; ZAKHAROV, Ye.I., inzh.; DEVYATKIN, V.I., inzh.;

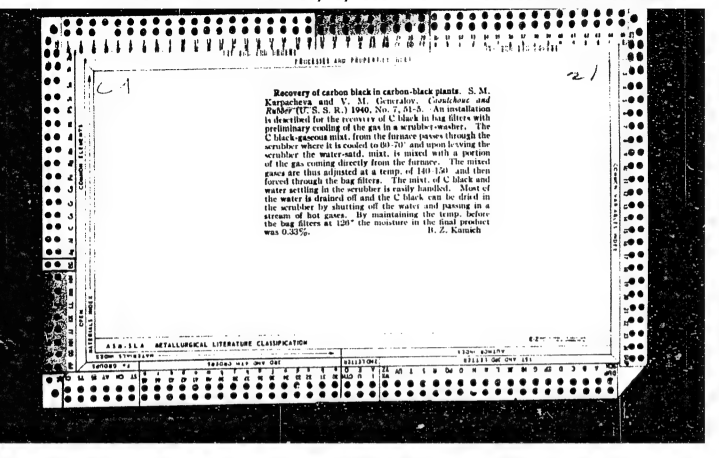
ZHDANOV, B.V., inzh.

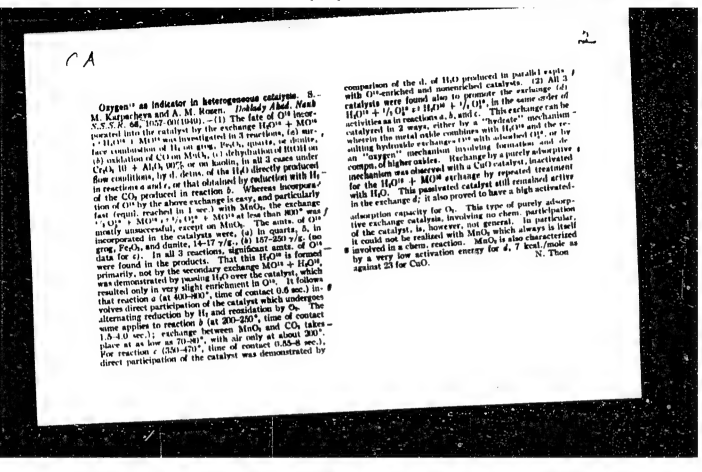
Study of the operation of a pulsating extraction sieve plate column. Khim. i neft. mashinostr. no.1:24-27 Ja 165.

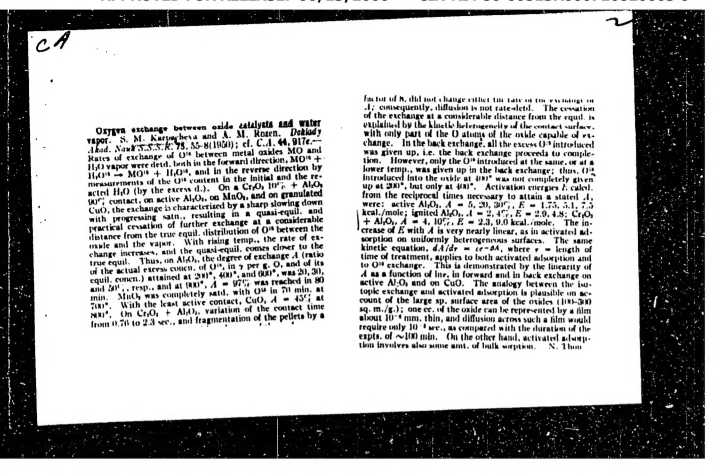
(MIRA 18:3)











KARPACHEVA, S.	through 2 stages: 1st there was a decompon of the alc as a result of the formation of alcoholates; then a decompon of the alcoholates back into the original products (alc and Al <sub>2</sub> O <sub>3</sub> ). Similar expts at 300 were carried out over kaolin, chromaluminum and zinc oxide. Dehydration resulted from the 1st 2, and dehydrogenation from the 3d. It is certain that all catalysts used form intermediate compds with alc, whether there is dehydration or not.	In previous work it was shown that the 018 in active oxides is capable of exchange with water vapor and certain liquids. Oh was introduced into the catalyst (Al203). Alcoholic vapors 1st at 2000, and then at 2000 were passed over the catalyst. The oxygen was exchanged from the catalyst to the alcoholic vapors 2-3 times faster than in the case of water. At the lower temp the exchange passed 214719	USER/Chemistry - Catalysts 21 Nov 51 "Oxygen Exchange Between Alcohol Vepors and Dehydration Catalysts," S. Karpacheva, A. Rozen "Dok Ak Mank SSER" Vol LXXXI, No 3, pp 425, 426	
		等場件等必要		,

